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### Title

Whole-Genome Analysis of Methyl tert-Butyl Ether-Degrading Beta-Proteobacterium  
Methylibium  
petroleiphilum PM1

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1       **Title:** Whole-Genome Analysis of Methyl *tert*-Butyl Ether (MTBE)-  
2       Degrading Beta-Proteobacterium *Methylibium petroleiphilum* PM1

3  
4       **Running Title:** Genome Sequence Analysis of *Methylibium petroleiphilum* PM1

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6

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1 **Abstract:**

2 *Methylibium petroleiphilum* PM1 is a methylotroph distinguished by its ability to  
3 completely metabolize the fuel oxygenate methyl *tert*-butyl ether (MTBE). Strain PM1  
4 also degrades aromatic (benzene, toluene, xylene) and straight chain (C<sub>5</sub>-C<sub>12</sub>)  
5 hydrocarbons present in petroleum products. Whole genome analysis of PM1 reveals a  
6 ~4-Mb circular chromosome and ~600-kb megaplasmid containing 3831 and 646 genes,  
7 respectively. Aromatic hydrocarbon and alkane degradation, metal resistance, and  
8 methylotrophy are encoded on the chromosome. The megaplasmid contains an unusual t-  
9 RNA island, numerous insertion sequences and large repeated elements including a 40-kb  
10 region also present on the chromosome and a 29-kb tandem repeat encoding phosphonate  
11 transport and cobalamin biosynthesis. The megaplasmid also codes for alkane  
12 degradation and was shown to play an essential role in MTBE degradation through  
13 plasmid curing experiments. Discrepancies between the IS element distribution pattern,  
14 the distribution of best BLASTP hits among major phylogenetic groups, and G+C content  
15 of the chromosome (69.2%) and plasmid (66%) together with comparative genome  
16 hybridization experiments suggest the plasmid was recently acquired and apparently  
17 carries the genetic information responsible for PM1's ability to degrade MTBE.  
18 Comparative genomic hybridization analysis with two PM1-like MTBE-degrading  
19 environmental isolates (~99% identical 16S rDNA sequences) showed that this plasmid  
20 was highly conserved (ca. 99% identical), whereas, the chromosomes were too diverse to  
21 conduct resequencing analysis. PM1's genome sequence provides a foundation to  
22 investigate MTBE biodegradation and explore genetic regulation of multiple  
23 biodegradation pathways in *M. petroleiphilum* and other MTBE-degrading beta-  
24 proteobacteria.

25

26

1 *Methylibium petroleiphilum* strain PM1, a newly described genus and species (57), is a  
2 motile bacterium belonging to the Comamonadaceae family of the beta-Proteobacteria  
3 and an important member of subsurface microbial communities in many gasoline-  
4 contaminated aquifers. Furthermore, PM1 is a methylotroph that can grow aerobically on  
5 the fuel oxygenate methyl *tert*-butyl ether (MTBE) and oxidize it completely to carbon  
6 dioxide (9, 35). MTBE is a suspected carcinogen that has contaminated drinking water  
7 wells throughout the US due to the preponderance of underground leaking storage tanks,  
8 the widespread usage of MTBE and its recalcitrance and mobility in groundwater. PM1  
9 can also oxidize aromatic hydrocarbons (toluene, benzene, o-xylene, and phenol) (20)  
10 and *n*-alkanes (C<sub>5</sub>-C<sub>12</sub>) (57; K. Hristova, unpublished data), and has been used in two  
11 bioaugmentation field trials in gasoline-contaminated aquifers in California (71) and  
12 Montana (18, 73). In contaminated sites amended with oxygen, *in situ* MTBE  
13 degradation was observed and corresponded to increases in native populations of  
14 *Methylibium* sp. (~99% similarity to PM1 based on 16S rDNA) (40, 71, 81). PM1-like  
15 bacteria occur naturally in a number of MTBE-contaminated aquifers in the US (47, 48,  
16 82), Mexico (21) and Europe (55, 61), and their presence has been correlated with MTBE  
17 degradation activity in numerous sites (48, 71, 82) using real-time PCR analysis (38).  
18 These results suggest that PM1-like organisms may play a major role in MTBE  
19 biodegradation under aerobic conditions in contaminated aquifers. The genetic basis for  
20 MTBE metabolism is not currently understood although there is general agreement that  
21 the initial enzymatic steps are similar to co-metabolic degradation pathways (27, 68, 74),  
22 and recent reports have described genes involved in degradation of MTBE downstream  
23 metabolites, 2-methyl-1,2-propanediol (29) and 2-hydroxyisobutyrate (61). The complex  
24 regulation of the metabolism of fuel hydrocarbons and MTBE, often occurring in  
25 mixtures, is relatively unknown, while limited studies showed that MTBE degradation  
26 could be inhibited in mixtures with BTEX compounds (20, 48).

27 In this paper, we present our analysis of the whole genome sequence of *M.*  
28 *petroleiphilum* PM1. We present comparative sequence analysis results between PM1  
29 and bacteria with homologous individual genes and operons as well as comparative  
30 whole genomic hybridization analysis between PM1 and PM1-like MTBE-degrading  
31 isolates (~99% identical 16S rDNA sequences) from gasoline-contaminated sites.

1 General genome features are discussed including interesting repeated elements, as well as  
2 genes and operons involved in methylotrophy, degradation of aromatic hydrocarbons,  
3 degradation of cyclic and straight-chain alkanes, cofactor biosynthesis, motility,  
4 secretion, and heavy metal resistance and transport. A noteworthy finding was the  
5 presence of a large ~600 bp plasmid in PM1 that was highly conserved among PM1-like  
6 bacteria. Furthermore, plasmid-curing experiments showed that the plasmid was essential  
7 for MTBE and TBA biodegradation in PM1. The PM1 genome sequence has provided a  
8 foundation for understanding novel pathways and interactions in this important  
9 subsurface bacterium as well as those in phylogenetically similar MTBE-degrading  
10 bacteria.

11

## 12 **MATERIALS AND METHODS**

13

### 14 **Bacterial strains used in genome sequence and comparative hybridization analyses.**

15 *Methylibium petroleiphilum* strain PM1 was used for whole genome sequencing at the  
16 Joint Genome Institute (Walnut Creek, CA). Strain PM1 was isolated from a sewage  
17 treatment plant biofilter used for treating discharge from oil refineries (34,9). Two  
18 MTBE-degrading bacterial pure cultures (MG4 and 312) were obtained from two  
19 different gasoline-contaminated aquifers in Northern California (47; Travis Air Force  
20 Base, Travis, CA and San Mateo, CA, respectively). Enrichment culturing was  
21 conducted in 10 mg/L MTBE mineral salts media (MSM; 56) with shaking at 150 rpm at  
22 room temperature. Enrichment cultures were plated onto 0.1X trypticase soy agar (TSA),  
23 and individual colonies were picked and grown in MSM with 10 mg/L MTBE and  
24 analyzed for MTBE degradation activity using purge-and-trap gas chromatography/mass  
25 spectrometry with reference to d<sub>12</sub>-MTBE as an internal standard (47). Culture purity  
26 was tested by plating (0.1X TSA) and 16S rDNA sequence analysis of colony genomic  
27 DNA.

28

29 **Sequencing, gene prediction and annotation.** Genomic DNA was isolated and purified  
30 from *M. petroleiphilum* PM1 and whole genome shotgun libraries (3-kb, 8-kb, and 40-kb

1 DNA inserts) were constructed and sequenced as previously described (14). After quality  
2 control of the 90,327 total initial reads of draft sequence, 83,180 sequences were  
3 assembled, producing an average of 10.7-fold coverage across the genome. The whole  
4 genome sequence was assembled using the Phred/Phrap/Consed package (P. Green,  
5 University of Washington) (25, 26, 33). The reads were assembled into 24 high-quality  
6 draft sequence contigs, which were linked into 3 larger scaffolds by paired-end sequence  
7 information. Gaps in the sequence were closed by either walking on gap-spanning clones  
8 or with PCR products generated from genomic DNA. Physical (un-captured) gaps were  
9 closed by combinatorial (multiplex) PCR. Sequence finishing and polishing added 308  
10 reads, and the final assessment of the genome assembly was completed as described  
11 previously (14). The final genome assembly quality of PM1 adheres to conventional  
12 standards of less than one error per 10000 bp. Each base is covered by at least 2 quality  
13 sequences, with an average of 10.7 fold coverage. Proper assembly was verified by  
14 fosmid coverage coupled with PCR data. Gene modeling and genome annotation was  
15 performed as previously described (14) to identify open reading frames likely encoding  
16 proteins (coding sequences [CDS]).

17

18 **Nucleotide sequence annotation and accession number.** The annotation is available on  
19 the Joint Genome Institute web-site (<http://genome.ornl.gov/microbial/rgel/>) and has been  
20 deposited in the GenBank/EMBL database under accession number  
21 NZ\_AAEM00000000.

22

23 **Comparative genomics.** Orthologs and CDSs unique to *M. petroleiphilum* PM1 were  
24 identified using the Integrated Microbial Genomes (IMG) system from the Joint Genome  
25 Institute. Results were based on BLASTP analysis with cutoff values of  $E < 10^{-5}$  and  
26 30% identity for orthologs and  $E < 10^{-2}$  and 20% identity for unique CDSs.

27

28 **Phylogenetic tree analysis.** Homologs of *M. petroleiphilum* PM1 translated CDSs were  
29 identified using BLASTP searches against the non-redundant (NR) GenBank database  
30 from National Center for Biotechnology Information. Sequences were aligned and  
31 alignments were refined using ClustalX version 1.8 (42) along with manual adjustments.

1 The *protdist* program and the *neighbor* program of the Phylip package (28) were used to  
2 generate the phylogenetic tree for MpeA3393. The pairwise parameters included gap  
3 opening = 35 and gap extension = 0.75. The multiple alignment parameters included gap  
4 opening = 15, gap extension = 0.3, delay divergence = 30%, and Gonnet series for the  
5 protein weight matrix.

## 7 **Comparative Genomic Hybridization and Comparative Genomic Sequencing**

8 **analyses.** Comparative Genomic Hybridization (CGH) was conducted in order to analyze  
9 conservation of genes from MTBE-degrading isolates MG4 and 312 with PM1 across the  
10 entire genome. High-density arrays (~400,000 oligomers) were designed and produced by  
11 NimbleGen Systems, Inc. (Madison, WI) using 29-mer probes every 26 bp for both  
12 strands of the entire genome and every 7 bases for both strands of the plasmid. Arrays  
13 were hybridized with labeled genomic preparations of MG4, 312 and PM1. Genomic  
14 DNA was isolated (5), and digested (5 µg per array) with 0.005 U DNase I (Amersham)  
15 in 1X One-Phor-All buffer (Amersham, Piscataway, NJ) at 37°C for 5 min with  
16 subsequent inactivation (95°C for 15 min). To the DNA digest were added 4 µL 5X  
17 Terminal Transferase Buffer, 1 nmol Biotin-N6-ddATP, and 25 U Terminal Transferase.  
18 The sample was incubated at 37°C for 90 min followed by inactivation at 95°C for 15  
19 min. Hybridization of arrays was conducted in 1 X hybridization buffer for 16 hr at 45°C  
20 using a Hybriwheel device (NimbleGen). PM1 was used as a reference in the analysis  
21 and was hybridized to separate arrays. Duplicate arrays were processed per strain. Arrays  
22 were washed with non-stringent wash buffer (6X SSPE, 0.01% [v/v] Tween-20) followed  
23 by two 5 min washes with stringent buffer (100 mM MES, 0.1 M NaCl, 0.01% [v/v]  
24 Tween-20) at 47.5°C. Arrays were stained with Cy3-streptavidin conjugate (Amersham  
25 Piscataway, NJ) for 10 min followed by washing in nonstringent buffer. Signal  
26 amplification was achieved by secondary labeling with biotinylated goat anti-streptavidin  
27 (Vector Laboratories, Burlingame CA), washing in nonstringent buffer and restaining  
28 with Cy3-streptavidin. Finally, arrays were washed in non-stringent wash buffer, in 0.5  
29 X SSC two times for 30 sec and in 70% ethanol for 15 sec. Arrays were spun dry by  
30 centrifugation. Scanning was conducted at 5-µm resolution with a Genepix 4000b  
31 scanner (Axon Instruments, Union City CA), and NimbleScan software (NimbleGen) was

1 used to obtain pixel intensities. For higher resolution resequencing of the MG4 and 312  
2 plasmids, arrays were synthesized and hybridized with genomic DNA from each strain  
3 and scanned as above. Single nucleotide polymorphism (SNP) positions were tested for  
4 uniqueness in the genome using custom algorithms (NimbleGen). The PM1 annotation  
5 (<http://genome.ornl.gov/microbial/rgel/>) was used to generate the output file in  
6 SignalMap analysis software (NimbleGen). The predicted SNPs were confirmed by  
7 producing and sequencing amplicons using PCR primers located external to the SNP  
8 location.

9

10 **Random mutagenesis, mutant characterization and plasmid curing of PM1.** In order  
11 to label the megaplasmid with a selectable marker, random transposon mutagenesis was  
12 employed using the mini transposon derivative, pTnMod-SmO containing the  
13 streptomycin/spectinomycin adenylyltransferase gene (*aadA*) and an oriR origin of  
14 replication between the inverted repeats (22). Electrocompetent PM1 cells were prepared  
15 by culturing in 0.5X Tryptic Soy Broth (TSB) at 27°C with shaking to log phase. Cells  
16 were collected by centrifugation, washed in 10% glycerol four times, and suspended in  
17 10% glycerol to a final volume of 100  $\mu$ l. A mixture of 50  $\mu$ l cells and 2  $\mu$ l pTnMod-  
18 SmO DNA (1  $\mu$ g/ $\mu$ l) was electroporated in 0.1 mm gap cuvettes using 1.8 kV, 200 Ohms,  
19 and 25  $\mu$ F capacitance settings (22) in a BioRad Gene Pulser Electroporator (BioRad,  
20 Hercules, CA). Following a 4 h recovery in 0.5X TSB at 27°C with shaking, transposon  
21 mutants were selected on 0.5X TSA plates with 50  $\mu$ g/ml streptomycin (Sm). Sm-  
22 resistant colonies were present after incubation for 5-6 days at 27°C and stable  
23 transposon integration was confirmed by PCR analysis of genomic DNA using pTnMod-  
24 SmO specific primers.

25 Using the rapid cloning strategy outlined by Dennis et al. (22), the <SmO> insert  
26 location was mapped in several PM1 subclones containing the oriR within the  
27 transposon. Briefly genomic DNA was extracted, digested with *Ava* II restriction  
28 endonuclease, self-ligated, and transformed into *E. coli* TOP10 cells (Invitrogen,  
29 Carlsbad, CA). The resulting transformants were selected on LB agar containing 50  
30  $\mu$ g/ml Sm. Sequencing with primers against the ends of the <SmO> insert was used to  
31 determine the exact insert location. One transposon-mutant MP0005 was shown to have



1 the <SmO> insert on the megaplasmid (in MpeB636). MP0005 was subjected to plasmid  
2 curing by heat stress as described by Trevors (78). Specifically, strain MP0005 was  
3 incubated at 37°C for 6-8 h before plating on 0.5X TSA. Following replica plating on  
4 0.5X TSA with and without 50 µg/ml Sm, Sm-sensitive colonies were selected and  
5 megaplasmid loss was confirmed by PCR analysis. MTBE and TBA degradation activity  
6 by strain MP0005 and a megaplasmid-free strain MP0007 were determined by gas  
7 chromatography analysis as previously described (35).

## 8 **Results and Discussion**

9 **General genome features of chromosome and megaplasmid.** The *Methylibium*  
10 *petroleiphilum* strain PM1 genome consists of a circular chromosome of 4,044,225 bp  
11 (Figure 1a.), and a megaplasmid (pPM1) of 599,444 bp (Figure 1b.) (Table 1). The  
12 genome encodes 4,477 putative CDSs, of which 964 are unique to PM1 based on  
13 BLASTP searches against NR. The pPM1-encoded proteins account for a  
14 disproportionately large number (382) of these unique genes. Of the remaining proteins,  
15 2801 could be assigned a putative function based on the KEGG (Kyoto Encyclopedia of  
16 Genes and Genomes) database. Analysis of the top BLAST hits (against completed  
17 genomes in KEGG) revealed the closest homolog was most often found in other beta-  
18 proteobacterial genomes (2332), with the most hits (790) to *Ralstonia solanacearum*  
19 followed by *Burkholderia pseudomallei* (497) and *Azoarcus* sp. EbN1 (413). This  
20 distribution appears to reflect that of the chromosome: 2210, 589 and 364 to beta-  
21 gamma-, and alpha-proteobacteria, respectively (Table 1). Interestingly, in contrast to the  
22 chromosome where beta- and gamma-proteobacteria account for 57.7% and 15.4% of its  
23 top BLAST hits respectively, the distribution of top hits between beta- (18.9%) and  
24 gamma- (15.6%) proteobacteria is nearly equivalent on the megaplasmid. The lower  
25 fraction of beta-proteobacteria-like CDSs in the megaplasmid is balanced by the large  
26 proportion of CDSs with no hits to KEGG genomes (47.7%) compared with the CDSs on  
27 the chromosome (9.9%). This surprising difference in the phylogenetic distribution of  
28 best hits together with the discrepancy in G+C content between the plasmid (66%) and  
29 the chromosome (69.2%) points to the likelihood that the plasmid was horizontally

1 acquired; further evidence for this statement is provided by conservation of the  
2 megaplasmid in other phylogenetically similar MTBE-degrading bacteria (discussed in  
3 detail later). Analysis of Clusters of Orthologous Genes (COG) distribution (77) showed  
4 that the most abundant groups (excluding no COG or general function) were amino acid  
5 transport and metabolism (7.0%), energy production (6.4%), and transcription (6.3%) on  
6 the chromosome, and replication, recombination and repair (8.0%), coenzyme transport  
7 and metabolism (7.0%), and inorganic ion transport and metabolism (5.3%) on the  
8 plasmid.

9       The chromosome contains a single ribosomal *rrn* operon (16S-tRNA<sup>ala</sup>-tRNA<sup>ile</sup>-  
10 23S-5S) and all genes coding for ribosomal proteins. Structural RNA genes for SRP  
11 RNA, *rnpB*, and tmRNA were present. Forty-two tRNA genes, evenly distributed on the  
12 chromosome (with the exception of a few clusters of 2 or 3 tRNAs), correspond to 40  
13 tRNA acceptors and can recognize all possible codons. A very unusual feature of pPM1  
14 is that it contains a single large cluster of 27 additional tRNA genes (25 are redundant  
15 with those on the chromosome, the two others do not have clear anticodons). This is the  
16 first report of such a large tRNA gene island, and the first report of such a cluster on a  
17 plasmid. The role of this island in translation, in genome evolution, or in positive  
18 selection of the plasmid in this or other bacterial strains is unclear.

19

20 **Cell motility, secretion and transport systems.** *M. petroleiphilum* PM1 possesses the  
21 genes necessary for flagellar biosynthesis (for one polar flagellum), chemotactic  
22 response, type IV pili synthesis, the type II secretion pathway as well as several genes  
23 related to the *Agrobacterium tumefaciens* type IV secretion pathway (Figure 2;  
24 Supplemental Table 1). Type IV secretion mechanisms are often involved in  
25 pathogenesis. However, homologs to only three of the five core type IV secretion  
26 proteins (VirB9, 10, 11, not VirB4 or 7) (6) were identified, so it is unclear at present if  
27 PM1 possesses a functional type IV secretion pathway. PM1 likely moves both by  
28 flagellar-facilitated swimming and pili-facilitated twitching motility. Three copies of *tra*  
29 genes on pPM1 suggest that PM1 may be capable of conjugative transfer, a possibility  
30 currently under investigation. Thirteen chromosomal and one plasmid born methyl-  
31 accepting chemotaxis proteins (MCP's) allow PM1 to respond to a range of

1 environmental stimuli (Supplemental Table 1). As in other organisms (83), MCP's in  
2 PM1 are found scattered throughout the genome. Only three MCP's are found within  
3 taxis operons – the *pilG-L* operon, the *cheYA(MCP)W* operon and the *flg/flh* gene cluster.  
4 The apparent mobility of MCP's, together with the fact that six PM1 MCP's appear to be  
5 paralogs, complicate function assignation. Nevertheless, there are five other MCP's in  
6 addition to those already mentioned whose gene environment may offer insight into their  
7 possible functions: two paralogous MCP's are located immediately downstream of and  
8 may be part of the same operon as the two toluene/benzene monooxygenase pathways;  
9 one of the *aer*-like MCP's is immediately downstream of, and in the same orientation as  
10 the LysR-type activator of the ribulose 1,5-bisphosphate carboxylase/oxygenase  
11 (RuBisCO) operon, which is upstream of the two regulatory genes; one MCP may be co-  
12 transcribed with a gene showing similarity to the direct oxygen sensor *dos* of *E. coli*; and  
13 one MCP may be co-transcribed with a gene showing low percent similarity to  
14 bacteriophytochromes and motility sensors. Neighbor-joining analysis of the putative  
15 PM1 MCP's against their homologs showed that eight MCP's cluster close to MCP1-4 of  
16 *E. coli*/ MCPA-D of *S. typhimurium*, two appear related to the aerotaxis and energy  
17 sensor AER of *E. coli*, and one is very similar to the twitching motility protein PilJ of *P.*  
18 *aeruginosa*.

19 Strain PM1 has two sets of genes coding for form I RuBisCO, *cbbL* (*mpeA1478*  
20 and *mpeA2782*) and *cbbS* (*mpeA1479* and *mpeA2783*) and associated enzymes required  
21 for CO<sub>2</sub> fixation via the Calvin cycle (Supplemental Table 1); however, this activity has  
22 not been demonstrated for PM1. A thorough search of the PM1 genome revealed the  
23 absence of key enzymes from each of the three other known CO<sub>2</sub> fixation pathways: 2-  
24 oxoglutarate:ferredoxin oxidoreductase and ATP citrate lyase (reductive TCA cycle); the  
25 acetyl-CoA synthase/CO dehydrogenase (reductive acetyl-CoA pathway); malonyl-CoA  
26 reductase and propionyl-CoA synthase (3-hydroxypropionate cycle) (4, 41). This strain  
27 possesses several ABC transporters for transport of inorganic ions such as nitrate, sulfate,  
28 magnesium, potassium, phosphate, phosphonate, as well as amino acids, branched chain  
29 amino acids, carbohydrates, long chain fatty acids, dipeptides/oligopeptides, polyamines,  
30 and antibiotics (Figure 2; Supplemental Table 2). In addition, putative

1 regulatory/signaling proteins, and cytochromes (based on CXXCH motifs) have been  
2 identified (Supplemental Table 3).

3

4 **Repeated elements.** The genome has a number of complex repetitive elements, including  
5 eight families of insertion sequences (ISmp1-8) (up to 12 copies) and two large genomic  
6 segments (29 and 40 kb) flanked by IS elements that have undergone what appears to be  
7 recent duplications (Figure 1). The two replicons do not equally share the repeated  
8 insertion elements; five of the eight families are located only on the chromosome and one  
9 family is strictly found on the plasmid. The distribution patterns of the IS elements, lends  
10 support to the dissimilar phylogenetic distribution of best KEGG hits among sequenced  
11 genomes and strengthens the notion of the megaplasmid's recent acquisition.

12         Parallel copies of ISmp8 flank two tandem copies of a 29-kb repeat, each  
13 consisting of two operons involved in phosphonate and cobalamin metabolism. The  
14 phosphonate operons (*PhnFDC-HtxFGHIJKLMN*) include putative C-P lyase subunits  
15 54-83% similar to those of *Pseudomonas stutzeri* WM88 (81) (Supplemental Table 2).  
16 The Htx and Phn C-P lyases support growth on methylphosphonate or additional  
17 alkylphosphonates, respectively; growth on these substrates is not yet known for PM1.  
18 Also contained in the repeat are cobalamin (vitamin B<sub>12</sub>) synthesis genes encoding the  
19 conversion of uroporphyrinogen III to cobinamide and the synthesis of  
20 dimethylbenzimidazole (DMB) in the aerobic pathway for cobalamin biosynthesis  
21 (*mpeB437-453, B472-488*) (Supplemental Table 1). Downstream of the tandem repeat are  
22 the remaining genes (*mpeB509-522*) for the covalent linkage of DMB, cobinamide and a  
23 phosphoryl group to complete the cobalamin synthesis pathway. Genes coding for the  
24 anaerobic pathway of cobalamin biosynthesis (*cbi* genes) are also present in the *cob*  
25 clusters, however, a complete pathway is lacking. PM1 also lacks a *cobG* encoding the  
26 monooxygenase that converts precorrin-3 to precorrin-4 in the aerobic pathway, however  
27 one or more of the multiple copies of *cbiG* (*mpeB479, 480, 444, 445*) may code for a  
28 functional enzyme that performs this reaction without oxygen (60). Cobalt, and  
29 cobalamin (vitamin B<sub>12</sub>) have been shown to enhance PM1's ability to grow on and  
30 degrade MTBE and its primary metabolite, *tert*-butyl alcohol (TBA) (K. Hristova,  
31 unpublished data) so it is not surprising that multiple copies of genes involved in

1 cobalamin synthesis are present in PM1 including tandem repeats of *cob* and *cbi* genes.  
2 Recently, Rohwerder et al. (61) showed that cobalamin synthesis affected the growth rate  
3 on the MTBE metabolites, TBA and 2-hydroxyisobutyrate (2-HIBA) for a beta-  
4 proteobacterial MTBE-degrading strain that was phylogenetically similar to PM1 (95.6%  
5 identical based on 16S rDNA sequence). In this strain, cobalt or cobalamin was necessary  
6 for activity of an enzyme, isobutyryl-CoA mutase involved in metabolism of 2-HIBA  
7 (61), and 99% identical homologs to this two-component mutase are present in PM1  
8 (*mpeB538/541*). As mentioned, a relatively large percentage of predicted proteins on the  
9 plasmid (7.0%) belong to the COG category for coenzyme transport and metabolism. A  
10 cluster of ethanolamine utilization (*eut*) genes (*mpeB0499-502*) found between the  
11 tandem repeats and third cobalamin cluster on the plasmid encode putative proteins 48-  
12 85% similar to EutJEMN from the cobalamin-dependent ethanolamine utilization  
13 pathway of *S. typhimurium* (50). The latter two proteins are homologs of carboxysome  
14 shell proteins (CcmKL). While *S. typhimurium* also contains the ethanolamine lyase  
15 subunits and regulator (EutBC and EutR) in its *eut* operon, in PM1 these genes  
16 (*mpeA2417-8, mpeA2415*) are on the chromosome.

17 The largest (40 kb) repeated element is present on both the plasmid and  
18 chromosome, and encodes a putative PinR-like site-specific recombinase, a replicative  
19 DNA helicase, 2 putative spoJ-like transcriptional regulators or plasmid partitioning  
20 proteins, a spoIIIE-like DNA translocase, a tellurite resistance protein, and many  
21 hypothetical products. The presence of the repeat on both replicons suggests a recent  
22 duplicative transfer event. Though the types of genes found in this region suggest a  
23 plasmid origin, both copies interrupt similar but non-identical copies of *dcd* genes (dCTP  
24 deaminase), and the direction of duplicative transfer remains to be proven. Since this  
25 duplication, the 40-kb repeat on pPM1 has been interrupted by a transposase between  
26 genes *mpeB0184* and *mpeB0187*.

27

28 **Heavy metal tolerance and metal homeostasis.** One interesting outcome of the genome  
29 analysis is evidence of PM1's potential resistance to heavy metals suggesting promise in  
30 using the organism to treat sites containing mixed wastes. Arsenic extrusion in PM1 is  
31 probably mediated by *arsRBC* present in two copies on the chromosome that are 58-81%

1 similar to each other (*mpeA1581-1584*, *arsHBCR*; *mpeA2343-4*, *A2347*, *arsCB* and *arsR*).  
2 While some other bacteria have five-gene operons (*arsRDABC*) and use the ArsAB  
3 pump, PM1 probably extrudes arsenite by a carrier protein, ArsB alone energized by  
4 membrane potential (62), although resistance to arsenic oxyanions needs to be evaluated.  
5 ArsC encodes an arsenic reductase responsible for the transformation of As(V) into  
6 As(III) and ArsR is a transcriptional repressor that responds to As(III) and Sb(III) (62).  
7 The function of the fourth gene product, ArsH, found in several bacteria (*Yersinia*  
8 *enterocolitica*, *Acidithiobacillus ferrooxidans*, *Pseudomonas aeruginosa*, *P. putida*  
9 KT2440) still remains unclear (10, 63). Three chromosomal copies of *chrA* (*mpeA2204*,  
10 *mpeA2205* and *mpeA2526*), belonging to the CHR family of transporters, may mediate  
11 chromate resistance in PM1. One copy of *ChrA* (*mpeA2204*) is 63 and 61% similar to its  
12 homolog in *Dechloromonas aromatica* RCB and *P. putida* KT2440, respectively,  
13 although a homologous chromate reductase (*ChrR*; 43) was not evident in PM1.

14 Genome analysis revealed 15 copper resistance genes in a large cluster (~14.4 kb at  
15 positions 1760297 – 1775675) on the PM1 chromosome with structural analogy to the  
16 plasmid-mediated (pMOL30) copper resistance cluster *copOAIPRSFG* in *Ralstonia*  
17 *metallidurans* (54) (Supplemental Table 2). The *CopOAIP-CopRS* cluster in PM1 is  
18 likely involved in the efflux of periplasmic copper (analogous to *cop* system of *P.*  
19 *syringae*, *R. metallidurans*, and *R. solanacearum*, and the *cos* system of *E. coli* [13]),  
20 whereas, the efflux of cytoplasmic copper is mediated by a P1-ATPase, *CopF1F2*  
21 (analogous to *R. metallidurans* *CopF*) (54). The genome of PM1 also has a putative  
22 chemiosmotic antiporter efflux system similar to *CzcCBA* of *R. metallidurans*, conferring  
23 resistance to Cd, Zn and Co (54). In addition to *copF*, there are two other genes encoding  
24 putative metal-transporting P1-type ATPases, *mpeA2479* and *mpeA3535*. Additional  
25 proteins potentially involved in metal transport include *NikBCDE* for nickel (*mpeA3117-*  
26 *3120*), *CbiOQMK* for cobalt and nickel (*mpeA2799-2802*), and *ModABC* (*mpeA3707*,  
27 *mpeA3714*, *mpeA3715*) for molybdenum uptake (Supplemental Table 2).

28 Ferric iron has also been shown to enhance PM1's ability to grow on and degrade  
29 MTBE and TBA (K. Hristova, unpublished data) so it is likely that active transport of  
30 iron is of particular importance. As with other Gram-negative bacteria, PM1 acquires its  
31 iron supply via  $Fe^{3+}$ -siderophores. *Fep* genes, which function in the synthesis of

1 polypeptides required for uptake of ferric enterobactin were identified inside each of the  
2 four *cob* operons in PM1 (Supplemental Table 1). *Polaromonas* sp., *R. ferrireducens*, and  
3 *M. flagellatus* all have iron transport genes either within or in close proximity to their *cob*  
4 operons (data not shown). Minimally, FepABC are required for ferric enterobactin  
5 uptake. *MpeA2292* and *mpeA2605* have been annotated as *fepA*, coding for an outer  
6 membrane receptor for an iron siderophore, however it is possible that *btuB* genes located  
7 near the *febBDC* genes are also involved in iron assimilation. The TonB-dependent  
8 energy transduction complex (*tonB*, *exbB* or *tolQ*, and *exbB*; Supplemental Table 1.)  
9 coded on the chromosome likely provides the mechanism for active transport of iron  
10 siderophores and cobalamin across the outer membrane (8, 38). The PM1 genome  
11 encodes about 39 putative proteins involved in iron transport and homeostasis, which  
12 implies the importance of iron in its physiology.

13

14 **Methylotrophy.** Methylotrophic metabolism of PM1 is of great interest because  
15 formaldehyde and formate are common intermediates of both methanol and MTBE or  
16 TBA oxidation by PM1 and other degraders (10, 59). *M. petroleiphilum* PM1 is capable  
17 of aerobic growth on methanol, formate, and succinate. Unlike other methylotrophic beta-  
18 proteobacteria, PM1 grows on MTBE, toluene, benzene, ethylbenzene, and  
19 dihydroxybenzoates as sole carbon sources (57, 59). PM1 possesses genes for the serine  
20 cycle and methylotrophy scattered in several different clusters on its chromosome (Figure  
21 1; Table 2). The strain does not grow on methylamine (K. Hristova, unpublished data),  
22 lacks a gene encoding the methylamine dehydrogenase (MADH) large subunit, and likely  
23 lacks MADH activity. Despite the ability of PM1 to grow on methanol, its genome lacks  
24 true homologs of *mxoF* and *mxoI*, known genes coding for the methanol dehydrogenase  
25 (MeDH) large and small subunits, present in several methylotrophs known to date. PM1  
26 contains a MeDH-like cluster *XoxF-J* (*mpeA3393-5*) that is present in *Methylobacterium*  
27 *extorquens* AM1 (17), which also contains the true *mxoF* cluster. Comparative sequence  
28 analysis of the product of gene *mpeA3393* revealed high similarity to MxoF (74% to *M.*  
29 *extorquens* AM1) and the *XoxF* homolog present in several non-methylotrophs (77% to  
30 *Burkholderia fungorum*). Based on phylogenetic analysis, *MpeA3393* clusters with the  
31 MxoF homologs of unknown function from other methylotrophic and non-methylotrophic

1 *Rhizobia* and *Burkholderia* spp., while true MeDH large subunits (MxaF) cluster together  
2 and are distinct from the MxaF homologs (Figure 3).

3 A cytochrome c-555 (*mpeA3394*) 56% similar to the  $c_H$  cytochrome of *M.*  
4 *capsulatus* Bath (electron donor to the oxidase in methylotrophic bacteria [1]) is found  
5 adjacent to the *mpeA3394*. A putative MxaJ/XoxJ (*mpeA3395*) shows 54% similarity  
6 with XoxJ from *Paracoccus denitrificans* and 42% similarity with MxaJ from *M.*  
7 *capsulatus* Bath. Five genes (*mpeA3829*, 2585-8) are involved in the biosynthesis of  
8 pyrroloquinoline quinone (PQQ), a cofactor of MeDH as well as quinoprotein ethanol  
9 dehydrogenase. A cluster of genes required for MeDH synthesis, *mxalKCASR*  
10 (*mpeA3273-3278*) is also present. To date, none of the gene clusters containing the XoxF  
11 homolog have been shown to be involved in methanol oxidation. Therefore, it is possible  
12 that a new enzyme is responsible for this function in the beta-proteobacterium *M.*  
13 *petroleiphilum* PM1.

14 Three different formate dehydrogenases are present in the PM1 genome, with  
15 homologs to *M. extorquens* and *M. capsulatus* Bath. The function of the tungsten-  
16 dependent formate dehydrogenase *fdh1* (*mpeA0337-339*), NAD-linked formate  
17 dehydrogenase *fdh2* (*mpeA3708-12*), and cytochrome-linked formate dehydrogenase  
18 *fdh3* (*mpeA1170-71*, 1173) for energy generation during growth on  $C_1$  substrates or for  
19 MTBE oxidation needs to be further explored. *MpeA3377* coding for a putative ABC-  
20 type tungstate transport system permease links gene clusters *fdh1* and *fdh2* (Table 2). The  
21 *fdh2* genes in PM1 have the same gene arrangement and significant sequence identity  
22 (52-81%) to the NAD-dependent formate dehydrogenase cluster *fdsGBACD* of *Ralstonia*  
23 *eutropha* (58). Pathways involved in metabolism and detoxification of formaldehyde, a  
24 central intermediate of both methanol and MTBE degradation by PM1 and other strains  
25 (36,58), may also function in MTBE metabolism. PM1 has two pathways for  
26 formaldehyde oxidation to  $CO_2$ , an  $H_4MPT$ -linked metabolic module that includes an  
27 archaeal-like gene cluster and an  $H_4F$ -linked metabolic module. Recently, phylogenetic  
28 analysis of a subset of bacterial and archaeal  $H_4MPT$ -linked  $C_1$  transfer genes placed  
29 PM1 sequences with other described beta-proteobacteria (45).

30



1 **Fuel hydrocarbon degradation pathways.** PM1 contains an operon (*mpeA0814-0821*)  
2 likely encoding for conversion of benzene to phenol (and catechol), and toluene to  
3 methylphenol (and methylcatechol) that is 62-74% similar to the benzene  
4 monooxygenase pathway in *P. aeruginosa* J1104 (BmoA-D1) (49) and 50-71% similar to  
5 the toluene para-monooxygenase (TpMO) pathway in *Ralstonia pickettii* PKO1  
6 (TbuA1UBVA2CX) (76) (Table 3). A second operon (*mpeA2539-2547*) is 55-74%  
7 identical to the first operon, however, it likely does not yield a functional monooxygenase  
8 since the TbuA1 homolog is interrupted by a transposon insertion and the TbuC homolog  
9 is a pseudogene. Both operons have two-component response regulator-sensor histidine  
10 kinases upstream and divergently transcribed (*mpeA0811-812*; *mpeA2536-2537*) although  
11 *mpeA2537* may be truncated due to the transposon insertion. *MpeA821* encodes a putative  
12 TbuX (65% similar to that in PKO1), an outer membrane protein regulated by TbuT and  
13 involved in toluene uptake (44). The BMO pathway has been implicated in benzene and  
14 toluene degradation (49), as has the TpMO pathway (76). In addition to benzene and  
15 toluene, PM1 has been shown to degrade *o*-xylene (19), although the biochemical  
16 pathway has not been elucidated to date. It is likely that *m*- and *p*-xylene can also be  
17 metabolized via the toluene monooxygenase (TMO) pathway of PM1 as described for  
18 PKO1 and other bacteria (30).

19 *M. petroleiphilum* PM1 grows on phenol, and two distinct clusters of  
20 dimethylphenol (*dmp*)-like genes are present (*mpeA2265-67*, *2272-86*; *mpeA3305-13*,  
21 *3321-25*) although the latter lacks the key structural gene *dmpP* so may not yield a  
22 functional phenol hydroxylase (PH). Gene products from the first cluster  
23 *dmpRKL MNOPQBCDEHFGI* are 60-83% similar to those on pVI150 in *Pseudomonas*  
24 *sp.* strain CF600 (66), including a multi-component PH, catechol 2,3-dioxygenase and the  
25 meta-cleavage pathway for catechol (Table 3). The second operon has transposon  
26 insertions inside *dmpC* and adjacent to *dmpO*. The PH subunits for the two operons are  
27 44-69% similar. The DmpR homologs (*mpeA3310*, *A2286*) are similar to TbuT (69 and  
28 65%) and may regulate TMO, PH and the meta-cleavage genes, since TbuT was shown to  
29 regulate these genes in PKO1 via separate promoters (11). However these operons are  
30 located together in PKO1, whereas, they are quite distant in PM1. PM1 can grow on  
31 phenol, and based on the presence of a complete *dmp* operon, it can likely degrade

1 alkylphenols as well, although it is not clear whether PH is essential for methylphenol  
2 degradation (as described for *P. stutzeri* OX1 [12]) or whether the TMO alone is capable  
3 of converting toluene to methylcatechol (as described for strain PKO1 [30]).

4 PM1 has nine CDSs encoding putative proteins with varying similarity to  
5 cyclohexanone monooxygenases (CHMOs) sometimes referred to as Baeyer-Villiger-  
6 type MOs (*mpeB579*, *B607*, *B610*, *A393*, *A898*, *A1038*, *A1351*, *A2885*, and *A2915*).  
7 Their protein products may play a role in hydroxylation of either alicyclic, aliphatic or  
8 aryl ketones to form a corresponding ester, which can easily be hydrolyzed. Alicyclic  
9 hydrocarbons represent up to 12% (wt/wt) of total hydrocarbons in petroleum mixtures  
10 (American Petroleum Institute). Aryl ketones such as acetophenone can be produced  
11 directly from atmospheric breakdown of ethylbenzene (a major petroleum component) or  
12 following abiotic conversion of ethylbenzene to ethylphenol (3) and subsequent  
13 biological conversion to the ketone. The putative CHMO genes are scattered across the  
14 genome and are not present in operons with other genes coding for subsequent  
15 metabolism after the MO reaction (i.e., esterases, alcohol and aldehyde dehydrogenases).  
16 The CHMOs have a narrow substrate range, possibly explaining the number of different  
17 flavoprotein MOs in PM1 with varying levels of similarity with representatives from this  
18 class (Table 2); the putative CHMOs in PM1 were more similar to phenylacetone MO  
19 (46-67%; ref. 53) than 4-hydroxyacetophenone MO (43-53%; ref. 45). In PM1, the nine  
20 Baeyer-Villiger MOs have a putative NADP<sup>+</sup>-binding site that is 72-88% similar to the  
21 proposed site in a CHMO from *Acinetobacter* sp. strain NCIMB 9871 (16).

22 An alkane monooxygenase pathway on pPM1 may facilitate PM1's growth on *n*-  
23 alkanes. In addition, alkane monooxygenase (hydroxylase) has been proposed to play a  
24 role in cometabolic MTBE hydroxylation since acetylene, an inactivator of short-chain  
25 alkane monooxygenase, was shown to inhibit MTBE degradation (68). In PM1 the  
26 hydroxylase subunit, AlkB (*mpeB0606*) is 69% and 66% similar to that of *Alcanivorax*  
27 *borkumensis* AP1 (72) and *P. putida* PGo1 (contained on the OCT plasmid) (79)  
28 respectively, and contains all 8 of the conserved His residues observed in other integral  
29 membrane binuclear-iron hydrocarbon monooxygenases (34). Also present are two  
30 rubredoxin genes (*mpeB0603* and *mpeB0602*), whose products are 76 and 78% similar,  
31 respectively, to rubredoxin 3 and 4 in *Gordonia* sp. strain TF6 (32). The rubredoxin (Rd)

1 coded by *mpeB603* is an AlkG1-type Rd, whereas, that coded by *mpeB602* is an AlkG2-  
2 type Rd, based on the CXXCG motif criteria described by van Beilen et al. (80); Only  
3 AlkG2-type Rds were shown to be functional in electron transfer from the rubredoxin  
4 reductase to alkane hydroxylase. In addition, *mpeB0601* codes for an ATP-dependent  
5 transcriptional regulator 38% similar to AlkS from *A. borkumensis* SK2 (36). Separated  
6 from the putative *alkS* by three hypothetical genes is a rubredoxin reductase, *alkT*  
7 (*mpeB0597*) whose protein product is 49% similar to that in *Gordonia* sp. TF6. PM1  
8 does not appear to possess any long-chain alkane (>C<sub>13</sub>) oxidation pathways such as an  
9 alkane dioxygenase (64), P-450 monooxygenase (2) or two alkane hydroxylase  
10 complexes (AlkMa and AlkMb) similar to *Acinetobacter* sp. strain M-1 (75), although  
11 PM1's single AlkB is 54% similar to both AlkMa and AlkMb. The gene organization of  
12 the *alk* operon in PM1 is somewhat similar to that of *Gordonia* sp. TF6 (*alkB2G1G2T*),  
13 except a transposase (*mpeB605*) and putative esterase (*mpeB604*) are between *alkB* and  
14 *alkG1G2* and as mentioned, a putative *alkS* and three hypothetical genes (*mpeB600-598*)  
15 are located between *alkG1G2* and *alkT* in PM1. Homologs to AlkHJKL from *P. putida*  
16 GPO1 coding for aldehyde dehydrogenase, alcohol dehydrogenase, acyl CoA synthetase,  
17 and outer membrane protein (79), respectively, were not present on the plasmid, although  
18 the PM1 chromosome contains homologs to AlkH (*mpeA2324*, 47% similar), AlkJ  
19 (*mpeA3803*, 58% similar), AlkK (*mpeA1769*, 71% similar) and AlkL (*mpeA3010*, 49%  
20 similar).

21 In addition, the PM1 chromosome contains a putative propane monooxygenase  
22 pathway (*mpeA950-953*) whose predicted proteins are 41-64% identical to PrmABCD in  
23 *Gordonia* sp. TY-5, coding for the large hydroxylase subunit, the NADH-dependent  
24 acceptor oxidoreductase, the small hydroxylase subunit and the coupling protein,  
25 respectively. The Prm complex in strain TY-5 was shown to catalyze the subterminal  
26 oxidation of propane yielding 2-propanol (51), while propane oxidation in PM1 is  
27 currently under investigation. As for PrmA in *Gordonia* TY-5, a pair of conserved Glu-  
28 X-X-His sequences are present in the putative PrmA of PM1 at residues 138-141 and  
29 237-240. The presence of these sequences is consistent with other monooxygenases in  
30 the binuclear-iron oxygenase family including soluble methane monooxygenases (23, 67)  
31 suggesting PrmA in PM1 may catalyze hydroxylation of propane. Like the operon in

1 strain TY-5, a chaperone similar to GroEL was adjacent to the *prm* cluster in PM1  
2 (*mpeA954*). Finally, PM1 has homologs to strain TY-5 alcohol dehydrogenases, *adh1*  
3 (*mpeA936*) and *adh3* (*mpeA599*) that are 72 and 83% similar, respectively, that may  
4 facilitate 2-propanol degradation. The putative monooxygenases in PM1 are summarized  
5 in Supplemental Table 3 including methanesulfonate monooxygenase, *msmA* and  
6 alkanesulfonate monooxygenase *ssuD*, which are part of *msmABDCEFGHG* and  
7 *ssuAADCBC* operons, respectively. PM1 may not utilize methanesulfonate since its *msmA*  
8 lacks the sequence CXH-X<sub>26</sub>-CXXH unique to methanesulfonate utilizers (7). In general,  
9 PM1 possesses several homologous genes with other soil bacteria including *Gordonia*,  
10 *Alcinovorax*, and *Pseudomonas* spp. capable of biodegradation of petroleum  
11 hydrocarbons as well as xenobiotic and recalcitrant compounds such as phthalates.

12

13 **MTBE biodegradation.** Though MTBE is a recent anthropogenic contaminant (released  
14 within the last 15 years), various microorganisms can utilize the compound for carbon  
15 and energy under aerobic conditions (31, 61, 65, 74). *M. petroleiphilum* PM1 is the best  
16 characterized of the few bacterial pure cultures reported to grow on and completely  
17 degrade MTBE and its daughter product TBA (20, 31, 37, 65). The genetic basis for  
18 MTBE and TBA conversion is not known, although different classes of monooxygenases  
19 have been proposed to play a role in metabolism or co-metabolism of these compounds  
20 (31, 37, 52, 68, 74), including P450-monooxygenase and alkane monooxygenase  
21 (hydroxylase) systems, the latter shown to play a role in cometabolic degradation of  
22 MTBE by *P. putida* GPo1 (69) and possibly also by *P. mendocina* KR-1 (70). A known  
23 inducer of alkane hydroxylase, dicyclopropylketone, was also shown to induce MTBE  
24 conversion to TBA in GPo1 (69). As reported above, an alkane MO (AlkB) system was  
25 detected in the PM1 genome on the megaplasmid. The AlkB in PM1 is likely involved in  
26 MTBE hydroxylation based on similarity to other AlkB proteins in organisms shown to  
27 be involved in MTBE degradation. Whereas, the K<sub>s</sub> values for MTBE in *n*-alkane grown  
28 GPo1 was reported to be quite high (20-40 mM), the apparent half saturation constant for  
29 MTBE by PM1 was 88 μM, which is in the range of K<sub>s</sub> values for MTBE by butane-  
30 degrading bacteria (52). Unlike GPo1 and KR-1, PM1 further degrades TBA, ultimately  
31 producing CO<sub>2</sub> and biomass. The putative AlkB in PM1 is proposed to only oxidize

1 MTBE and not TBA based on kinetics experiments with MTBE- and TBA-grown cells  
2 (19). Two separate enzyme systems were also suggested for MTBE and TBA degradation  
3 by *Hydrogenophaga flava* ENV735 (37). Because of its potential role in MTBE  
4 metabolism, the coding region for AlkB is the focus of current gene knockout studies.  
5 Biodegradation of a similar molecule, ethyl-*tert* butyl ether (ETBE), occurs via a  
6 cytochrome P450 pathway in *Rhodococcus ruber* IFP2001 (15). However, homologs of  
7 protein complexes involved in ETBE degradation from *R. ruber* were not found in PM1;  
8 like GPo1 (69), PM1 has not been shown to degrade ETBE.

9 Many pollutant degradation genes are located on bacterial catabolic plasmids.  
10 Significantly, the two strains, MG4 and 312 that are capable of MTBE degradation had a  
11 nearly identical plasmid to that of PM1 (ca. 99% identical) as determined by comparative  
12 genomic sequencing analysis. The MG4 and 312 plasmids showed only 5 or 4 SNPs  
13 respectively relative to PM1 (Table 3). MG4 and 312 plasmids also lacked transposase  
14 genes (three copies of Tra5 and transposase-8 and one copy of a DDE-domain  
15 transposase) that were present on the PM1 plasmid and a 1.2 kb deletion putatively  
16 containing an esterase/lipase gene (*mpeB604*) and a DDE-domain transposase (*mpeB605*)  
17 (Table 3). The promoter and coding region for *alkB* (*mpeB606*) did not appear to be  
18 affected by this deletion since it is significantly upstream although there was a SNP  
19 mapped within *alkB* of MG4 and 312 resulting in a putative amino acid change. As  
20 mentioned, two other PM1 plasmid-encoded genes *mpeB541* and *mpeB538* code for  
21 putative large and small subunits of isobutyryl-coenzyme A (CoA) mutase, respectively.  
22 The plasmids of strains MG4 and 312 also contained identical copies of *mpeB541* and  
23 *mpeB538*. These gene products were shown to have 99% identical homologs in *Ideonella*  
24 sp. strain L108 predicted to allow conversion of 2-HIBA to 3-hydroxybutyrate in the  
25 presence of CoA and ATP (61). It is not known whether these mutase genes are  
26 contained on a megaplasmid in L108 although horizontal gene transfer is often evoked  
27 when such high similarities in gene sequences between bacteria are observed. In addition  
28 to *alkB*, the PM1 plasmid (as well as MG4 and 312 plasmids) contains a gene coding for  
29 3-hydroxybutyryl-CoA dehydrogenase (*mpeB0547*), putatively involved in conversion of  
30 3-hydroxybutyryl-CoA, a proposed MTBE-metabolite (61), to acetoacetyl-CoA.

1           The role of the megaplasmid in MTBE and TBA degradation was clearly  
2 demonstrated by curing experiments. Chemical analysis of MTBE and TBA degradation  
3 by the MP0005 parent strain and the MP0007 strain lacking the megaplasmid (as  
4 evidenced by PCR analysis and loss of Sm-resistance) showed that only the former was  
5 able to degrade MTBE and TBA (Figure 4). This result is consistent with our proposal  
6 that key genes involved in both MTBE and TBA degradation are located on the PM1  
7 megaplasmid. Since two different monooxygenases are proposed to be involved in  
8 MTBE and TBA degradation (19; K. Hristova, unpublished data), at least some of the  
9 required protein subunits for conversion of MTBE to TBA and conversion of TBA to the  
10 putative 2-methyl-1,2-propanediol are coded on the megaplasmid. In addition, the  
11 possible role of selected pPM1 proteins in MTBE/TBA oxidation, based on cDNA  
12 microarray results, is currently under investigation by gene knockout methods. It is  
13 noteworthy that the PM1 plasmid did not contain predicted proteins with significant  
14 homology to those proposed in degradation of 2-methyl-1,2-propanediol by  
15 *Mycobacterium austroafricanum* IFP 2012 (29), although the putative aldehyde  
16 dehydrogenase coded by *mpeA1909* on the chromosome was 54% similar to MpdC  
17 (hydroxyisobutyraldehyde dehydrogenase) and the alcohol dehydrogenase coded by  
18 *mpeA945* was 45% similar to MpdB (2-methyl-1,2-propanediol dehydrogenase). While  
19 the relevant alcohol and aldehyde dehydrogenases may be encoded on the chromosome,  
20 additional plasmid-encoded dehydrogenases may be more plausible and are currently  
21 being investigated for their putative role in the MTBE degradation pathway.

22

23 **Concluding remarks.** Prior to sequencing its genome, it was not known that PM1  
24 possessed a 600-kb megaplasmid, much less that the plasmid contained candidate CDSs  
25 coding for the MTBE and TBA monooxygenases and enzymes involved in downstream  
26 reactions. It is noteworthy that MTBE-degrading strains from diverse locations including  
27 a biofilter treating wastewater in Southern California (PM1) and two distinct aquifers in  
28 Northern California (MG4 and 312) possess a nearly identical plasmid. The presence of  
29 this highly conserved megaplasmid among PM1-like MTBE-degraders, along with its  
30 different G+C content, its unique IS complement and the unique phylogenetic  
31 distribution of its gene products, together raise interesting questions concerning

1 horizontal gene/plasmid transfer and evolution of pathways via plasmid-mediated  
2 mechanisms. With the whole genome sequence, putative aromatic hydrocarbon and  
3 alkane degradation pathways were also identified providing a basis to study the complex  
4 regulation of fuel hydrocarbon degradation in this novel subsurface bacterium; this is  
5 important since substrate interactions are expected to influence the success of  
6 bioremediation strategies for gasoline-contaminated sites. In addition to comparative  
7 genomics approaches, whole genome microarray and 2-D gel electrophoresis experiments  
8 are being conducted to identify genes and proteins unique to MTBE degradation. PM1  
9 can serve as a model for other MTBE-degrading methylotrophs such that the knowledge  
10 gained from analysis of its genome, transcriptome and proteome can be applied to PM1-  
11 like bacteria. An understanding of the MTBE degradation pathway and its regulation will  
12 allow for optimization of MTBE bioremediation and the ability to monitor this unique  
13 process *in situ* using molecular tools.

14

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1

**TABLE 1. General features of the *Methylibium petroleiphilum* PM1 genome<sup>1</sup>**

<b>Feature</b>	<b>Chromosome</b>	<b>Megaplasmid</b>
Total bases	4,044,225	599,444
CDS, total	3831	646
CDS, percent	92.4	86.3
Average CDS size, bp	1059	951
RNA operons (16S-23S-5S)	1	0
tRNAs	42	27
GC percent	69.20	65.98
Repeated IS elements	25	11
Alpha-proteobacterial KEGG hits	364 (9.5%)	31 (4.8%)
Beta-proteobacterial KEGG hits	2210 (57.7%)	122 (18.9%)
Gamma-proteobacterial KEGG hits	589 (15.4%)	101 (15.6%)
No hits in KEGG genome DB	378 (9.9%)	308 (47.7%)
Conserved hypothetical	684 (17.9%)	59 (9.1%)
Hypothetical (Unique)	582 (15.2%)	382 (59.1%)
Total proteins with EC assignment	682	

2 <sup>1</sup> The BLASTP threshold e-score was 1e-05.

3

1 **TABLE 2.** *M. petroleiphilum* PM1 genes putatively involved in metabolism of methanol.

<b>Gene ID</b>	<b>Gene Name</b>	<b>COG no.</b>	<b>Predicted Role</b>
<b>PQQ synthesis cluster</b>			
MpeA3829	<i>pqqA</i>		Pyrroloquinoline quinone biosynthesis protein A
MpeA2585	<i>pqqB</i>		Pyrroloquinoline quinone biosynthesis protein B
MpeA2586	<i>pqqC</i>	COG5424	Pyrroloquinoline quinone biosynthesis protein C
MpeA2587	<i>pqqD</i>		Pyrroloquinoline quinone biosynthesis protein D
MpeA2588	<i>pqqE</i>	COG0535	Pyrroloquinoline quinone biosynthesis protein E
<b>C1 transfer cluster 1</b>			
MpeA2606	<i>mptG</i>	COG1907	synthase, methanopterin biosynthesis
MpeA2607	<i>mptD</i>		methylene tetrahydromethanopterin dehydrogenase
MpeA2608		COG2232	ATP-dependent carboligase
MpeA2609	<i>mch</i>	COG3252	methenyl-H <sub>4</sub> MPT cyclohydrolase
MpeA2610		COG0189?	putative H <sub>4</sub> MPT biosynthesis protein
MpeA2611		COG1767	triphosphoribosyl-dephosphoCoA-synthetase
MpeA2612	<i>fae</i>	COG1795	formaldehyde-activating enzyme
MpeA2613	<i>folB</i>	COG1539	Dihydroneopterin aldolase
MpeA2614		COG3284	putative transcriptional activator
MpeA2615		COG1036	similar to Archaeal flavoprotein
MpeA2616	<i>folP</i>	COG0294	Dihydropteroate synthase
MpeA2617			involved in H <sub>4</sub> MPT -linked formaldehyde oxidation
MpeA2618		COG1891	uncharacterized protein in Archaea, involved in denovo synthesis of NAD in prokaryotes & eukaryotes
MpeA2619		COG2054	uncharacterized Archaeal kinase
MpeA2620		COG1548	putative transcriptional regulator/kinase
MpeA2621		COG1821	putative ATP utilizing enzyme
MpeA2622		COG1411	involved in H <sub>4</sub> MPT-linked formaldehyde oxidation
MpeA2623	<i>fhcB</i>	COG1029	formyltransferase/hydrolase complex, beta subunit
MpeA2624	<i>fhcA</i>	COG1229	formyltransferase/hydrolase complex, alpha subunit
MpeA2625	<i>fhcD</i>	COG2037	formyltransferase/hydrolase complex, delta subunit
MpeA2626	<i>fhcC</i>	COG2218	formyltransferase/hydrolase complex, gamma subunit
MpeA2627	<i>fae</i>	COG1795	putative formaldehyde-activating enzyme, present in Archaea
<b>Primary oxidation Methylamine DH cluster</b>			
MpeA2651		COG4977	Transcriptional regulator
MpeA2652	<i>cytC</i>	COG2010	Cytochrome C
MpeA2653	<i>mauA</i>		methylamine dehydrogenase small subunit
MpeA2654	<i>mauD</i>		methylamine utilization protein
<b>C1 transfer +C1 assimilation cluster 2</b>			
MpeA3251		COG1560	layroyl acyltransferase
MpeA3254	<i>mcl</i>		methyltrophic malyl-CoA lyase
MpeA3255	<i>ppc</i>	COG2352	methylotrophic phosphoenolpyruvate carboxylase
MpeA3256	<i>mtkB</i>	COG0074	malate thiokinase
MpeA3257	<i>mtkA</i>		malate thiokinase
MpeA3258	<i>fch</i>	COG3404	methenyl tetrahydrofolate cyclohydrolase
MpeA3259	<i>mtdA</i>	COG0300	methylene-H <sub>4</sub> MPT/ methylene-H <sub>4</sub> F dehydrogenase
MpeA3260			hydroxypyruvate reductase
MpeA3261	<i>sgaA</i>	COG0075	serine glyoxylate aminotransferase
MpeA3262	<i>glyA</i>	COG0112	serine hydroxymethyltransferase
MpeA3263	<i>gck</i>	COG2379	glycerate kinase
MpeA3264	<i>ftfL</i>	COG2759	formyl-tetrahydrofolate ligase
MpeA3265	<i>folC</i>	COG0285	dihydrofolate synthetase

1 **TABLE 2.** (continued)

<b>Gene ID</b>	<b>Gene Name</b>	<b>COG no.</b>	<b>Predicted Role</b>
<b>Primary oxidation - Mxa cluster</b>			
MpeA3273	<i>mxal</i>	COG1764	MxaL protein
MpeA3274	<i>mxak</i>		MxaK protein
MpeA3275	<i>mxac</i>	COG2425	MxaC protein
MpeA3276	<i>mxaa</i>		MxaA protein
MpeA3277	<i>mxas</i>	COG1721	MxaS protein
MpeA3278	<i>mxar</i>	COG0714	putative MxaR - involved in the regulation of formation of active MDH
MpeA3391		COG1521	Putative transcriptional regulator
MpeA3393	<i>mxaf?</i>	COG4993	methanol dehydrogenase large subunit homolog
MpeA3394	<i>cytC</i>	COG2010	cytochrome C-555
MpeA3395	<i>mxaj</i>		MxaJ like protein
<b>Primary oxidation - Formate dehydrogenase cluster</b>			
<b>NAD-linked molybdenum dependent formate dehydrogenase</b>			
MpeA3706		COG2998	ABC type tungstate transport system permease
MpeA3708	<i>fdh2A</i>	COG1905	formate dehydrogenase gamma subunit
MpeA3709	<i>fdh2B</i>	COG1894	formate dehydrogenase beta subunit
MpeA3710	<i>fdh2C</i>	COG3383	formate dehydrogenase alpha subunit
<b>Tungsten-containing formate dehydrogenase</b>			
MpeA0337	<i>fdh1A</i>	COG3383	formate dehydrogenase alpha subunit
MpeA0338	<i>fdh1B</i>	COG1894	formate dehydrogenase beta subunit
<b>Cytochrome-linked formate dehydrogenase</b>			
MpeA1170	<i>fdh3A</i>	COG0243	formate dehydrogenase alpha subunit
MpeA1171	<i>fdh3B</i>	COG0437	formate dehydrogenase beta subunit
MpeA1173	<i>fdh3C</i>	COG2864	formate dehydrogenase gamma subunit
<b>C1 assimilation</b>			
MpeA1014	<i>mcmB</i>	COG1884	Methylmalonyl-CoA mutase, beta subunit
MpeA1014	<i>mcmA</i>		Methylmalonyl-CoA mutase, alpha subunit
MpeA1015	<i>meaB</i>	COG1703	GTPase/ATPase, essential for methylmalonyl-CoA mutase reaction
MpeA1016	<i>pccB</i>		
MpeA1017	<i>pccA</i>	COG4770	Propionyl-CoA carboxylase, alpha subunit
MpeA1018	<i>epm</i>		Methylmalonyl-CoA epimerase
<b>Succinate dehydrogenase cluster</b>			
MpeA2165	<i>gltA</i>	COG0372	Citrate (Si)-synthase
MpeA2166		COG2938	
MpeA2167	<i>sdhB</i>	COG0479	Succinate dehydrogenase/fumarate reductase Fe-S protein subunit'
MpeA2168	<i>sdhA</i>	COG1053	Succinate dehydrogenase/fumarate reductase
MpeA2169	<i>sdhD</i>	COG2142	Transmembrane succinate dehydrogenase flavoprotein subunit
MpeA2170	<i>sdhC</i>	COG2009	Succinate dehydrogenase/fumarate reductase
MpeA2172	<i>mdh</i>	COG0039	malate dehydrogenase cytochrome b subunit hydrophobic membrane anchor subunit

2

3

1 **TABLE 3.** *M. petroleiphilum* PM1 genes putatively involved in metabolism of aromatic  
 2 hydrocarbons denoting the predicted role of the gene product and percent similarity with  
 3 well-characterized homologs.  
 4

Operon	Gene ID	Gene Name	Predicted Role	% Similarity with PKO1 <sup>1</sup>	
<i>tbu</i>					
<b>Operon I</b>	MpeA0811		two-component response regulator		
	MpeA0812		two-component sensor histidine kinase		
	MpeA0813		hypothetical		
	MpeA0814	<i>bmoA/tbuA1</i>	toluene monooxygenase alpha subunit	84	
	MpeA0815	<i>bmoB/tbuU</i>	toluene monooxygenase gamma subunit	81	
	MpeA0816	<i>bmoC/tbuB</i>	toluene monooxygenase ferredoxin	74	
	MpeA0817	<i>bmoD1/tbuV</i>	toluene monooxygenase activator	76	
	MpeA0818	<i>tbuA2</i>	toluene monooxygenase beta subunit	82	
	MpeA0819	<i>tbuC</i>	toluene monooxygenase oxidoreductase	64	
	MpeA0820		probable Zn-alcohol dehydrogenase		
MpeA0821	<i>tbuX</i>	toluene facilitator	65		
<i>tbu</i>					
<b>Operon II</b>	MpeA2536		two-component response regulator		
	MpeA2537		two-component sensor histidine kinase		
	MpeA2538	<i>tnp</i>			
	MpeA2539	<i>bmoA/tbuA1</i>	toluene monooxygenase alpha subunit C-terminal end	81	
	MpeA2540	<i>bmoB/tbuU</i>	toluene monooxygenase gamma subunit	76	
	MpeA2541	<i>bmoC/tbuB</i>	toluene monooxygenase ferredoxin	76	
	MpeA2542	<i>bmoD1/tbuV</i>	toluene monooxygenase activator	77	
	MpeA2543	<i>tbuA2</i>	toluene monooxygenase beta subunit	79	
	MpeA2544	<i>tbuC</i>	toluene monooxygenase oxidoreductase N-terminal end	69	
	MpeA2545	<i>tbuC</i>	toluene monooxygenase oxidoreductase C-terminal end	60	
					<b>% Similarity with CF600<sup>2</sup></b>
	<b>Operon I</b>	MpeA2286	<i>dmpR</i>	regulator (HylR family)	65
		MpeA2285	<i>dmpK</i>	phenol hydrolase assembly	60
		MpeA2284	<i>dmpL</i>	phenol hydrolase beta subunit	63
MpeA2283		<i>dmpM</i>	phenol hydrolase activator	76	
MpeA2282		<i>dmpN</i>	phenol hydrolase alpha subunit	77	
MpeA2281		<i>dmpO</i>	phenol hydrolase gamma subunit	63	
MpeA2280		<i>dmpP</i>	phenol hydrolase reductase	73	
MpeA2279		<i>aphT</i>	regulator (LysR family)		
MpeA2278		<i>dmpQ</i>	catechol-2,3-dioxygenase ferredoxin	62	
MpeA2277		<i>dmpB</i>	catechol-2,3-dioxygenase	70	
MpeA2276		<i>aphY</i>	conserved in some dmp-like operons		
MpeA2275		<i>dmpC</i>	2-hydroxymuconic semialdehyde dehydrogenase	83	
MpeA2274		<i>dmpD</i>	2-hydroxymuconic semialdehyde hydrolase	81	
MpeA2273		<i>dmpE</i>	2-hydroxypent-2,4-dienoate hydratase	78	
MpeA2272		<i>dmpH</i>	4-oxalocrotonate decarboxylase	81	
MpeA2267		<i>dmpF</i>	acetaldehyde dehydrogenase (acylating)	69	



	MpeA2266	<i>dmpG</i>	4-hydroxy-2-isovalerate aldolase	70
	MpeA2265	<i>dmpI</i>	4-oxalocrotonate isomerase	73
<b><i>dmp</i></b>				
<b>Operon II</b>	MpeA3328	<i>dmpC</i>	2-hydroxymuconic semialdehyde dehydrogenase N-terminal end	72
	MpeA3327	<i>tnp</i>	transposase	
	MpeA3326	<i>dmpC</i>	2-hydroxymuconic semialdehyde dehydrogenase C-terminal end	83
	MpeA3325	<i>dmpE</i>	2-hydroxypent-2,4-dienoate hydratase	73
	MpeA3324	<i>dmpH</i>	4-oxalocrotonate decarboxylase	72
	MpeA3323	<i>dmpI</i>	4-oxalocrotonate isomerase	66
	MpeA3322	<i>dmpF</i>	acetaldehyde dehydrogenase (acylating)	67
	MpeA3321	<i>dmpG</i>	4-hydroxy-2-isovalerate aldolase	73
	MpeA3313	<i>aphT</i>	regulator (LysR family)	
	MpeA3312	<i>dmpQ</i>	catechol-2,3-dioxygenase ferredoxin	63
	MpeA3311	<i>dmpB</i>	catechol-2,3-dioxygenase	68
	MpeA3310	<i>dmpR</i>	regulator (HylR family)	61
	MpeA3309	<i>dmpK</i>	phenol hydrolase assembly	64
	MpeA3308	<i>dmpL</i>	phenol hydrolase beta subunit	67
	MpeA3307	<i>dmpM</i>	phenol hydrolase activator	76
	MpeA3306	<i>dmpN</i>	phenol hydrolase alpha subunit	74
	MpeA3305	<i>dmpO</i>	phenol hydrolase gamma subunit	57
	MpeA3304	<i>tnp</i>	transposase	

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1

2 <sup>1</sup> Percent similarity with homologs in *Ralstonia pickettii* PKO1 (Tao et al., 2004; Accession nos.  
3 AY541701, AF100782, and U72645).

4 <sup>2</sup> Percent similarity with homologs in *Pseudomonas sp.* strain CF600 (Shingler et al., 1992;  
5 Accession nos. X60657, X60835, X52805, X33263, X60276, and X68033).

6

7

8

**TABLE 4. Genomic differences between the plasmid of PM1 with those from isolates MG4 and 312.**

Genome Location <sup>a</sup>	DNA Change <sup>b,c</sup>	Gene	pPM1	MG4	312
3921	Δ (1337 bp)	ISMca2 tnp tnp OCP	Not present	Present	Present
168369	Δ (1321 bp)	ISMca2 tnp tnp OCP	Not present	Present	Present
194159	A → G	Upstream of MpeB219	A	G	G
267528	T → G	Upstream of MpeB319	T	G	T
315627	T → G	Upstream of MpeB365	T	G	G
465830	C → G	<i>cobA</i> (Mpe515)	His176	Asp176	Asp176
504193	Δ (529 bp)	TIS 1021 tnp protein	Not present	Present	Present
536177	Δ (1370 bp)	ISMca2 tnp tnp OCP	Not present	Present	Present
556849	Δ (1229 bp)	MpeB604-B605 lipase/esterase, tnp-IS5 family	Not present	Present	Present
559059	T → G	<i>alkB</i> (MpeB606)	Phe264	Cys264	Cys264
Total SNPs				5	4
Missing genes				9	9

1 <sup>a</sup> Location in PM1 genome. SNP = single nucleotide polymorphism.

2 <sup>b</sup> Deletion (Δ) or SNP.

3 <sup>c</sup> Change relative to PM1 sequence. Predicted amino acid changes are noted where  
4 appropriate.

5

1 **Figure Legends**

2

3 **Figure 1.** Circular maps of the chromosome (a) and megaplasmid pPM1 (b) of *M.*  
 4 *petroleiphilum* PM1. The outer rings 1 and 2 show all CDSs colored by functional  
 5 category (dark gray = hypothetical proteins; light gray = conserved hypothetical and  
 6 unknown function; brown = general function prediction; red = replication and repair;  
 7 green = energy metabolism; blue = carbon and carbohydrate metabolism; cyan = lipid  
 8 metabolism; magenta = transcription; yellow = translation; orange = amino acid  
 9 metabolism; pink = metabolism of cofactors and vitamins; light red = purine and  
 10 pyrimidine metabolism; lavender = signal transduction; sky blue = cellular processes;  
 11 pale green = structural RNAs). Genes coding for major metabolic features (green, =  
 12 methylotrophy; red = aromatic hydrocarbon degradation; light blue = alkane degradation)  
 13 are shown on rings 3 and 4. Large repeat regions are indicated on rings 5 and 6. The  
 14 larger repeated regions colored light gray (29 kb repeat) and dark gray (40 kb repeat) are  
 15 described in the text, while the other colors represent repeated IS elements, ISmp1-8.  
 16 Ring 7 shows the deviation from the average G+C and the innermost ring 8 shows the  
 17 GC skew (G-C)/(G+C). The plasmid and chromosome are not drawn to scale.

18

19

20 **Figure 2.** Schematic diagram of a *M. petroleiphilum* PM1 cell showing structural features  
 21 and cellular processes including predicted methylotrophy and fuel hydrocarbon  
 22 degradation pathways. Two adjacent arrows imply multiple steps. Abbreviations: MTBE,  
 23 methyl *tert*-butyl ether; HMTBE, hydroxymethyl *tert*-butyl ether; TBF, *tert*-butyl  
 24 formate; TBA, *tert*-butyl alcohol; HIBA, hydroxyisobutyric acid. Systems involved in  
 25 metal resistance and transport, secretion, motility, chemotaxis and electron transport are  
 26 shown within or associated with the cell membrane. Cofactors are labeled green and  
 27 electron donors are labeled red in the cytosol. Figure is not drawn to scale.

28

29 **Figure 3.** The phylogenetic tree of MpeA3393, which putatively codes for the large  
 30 subunit of methanol dehydrogenase. The protein product from MpeA3393 is most  
 31 similar to glucose dehydrogenase [GluDH] from *B. fungorum* (81% similar; Acc. No.

1 ZP\_00283396) and methanol dehydrogenase large subunit [Mdh\_large] from *M.*  
2 *capsulatus* Bath (75% similar; Acc. No. AAU90462). Gene names are in square brackets.

3

4 **Figure 4.** MTBE and TBA degradation by *M. petroleiphilum* strain PM1 by the parent  
5 strain MP0005 that carries the <SmO> marker on the megaplasmid (MTBE - ●, TBA -  
6 ○) and the megaplasmid-free mutant MP0007 (MTBE - ■, TBA - □).

7

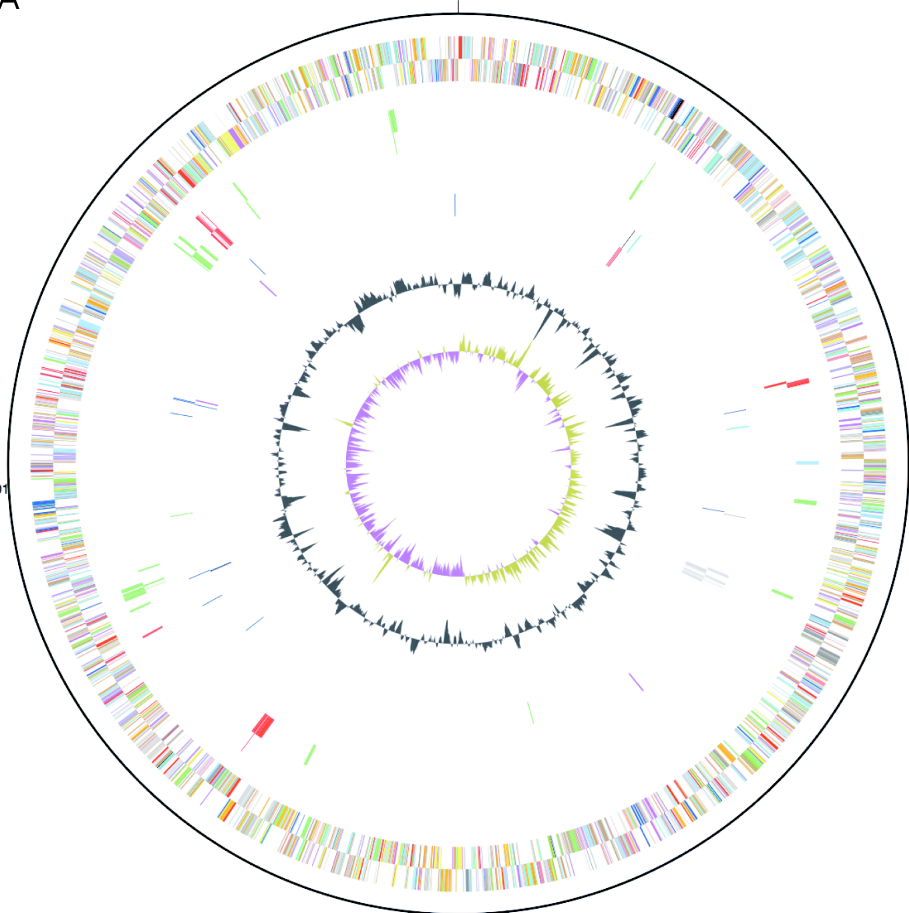
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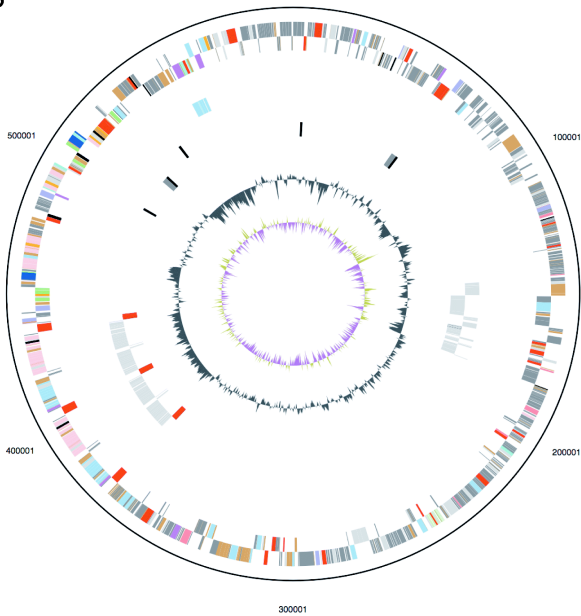
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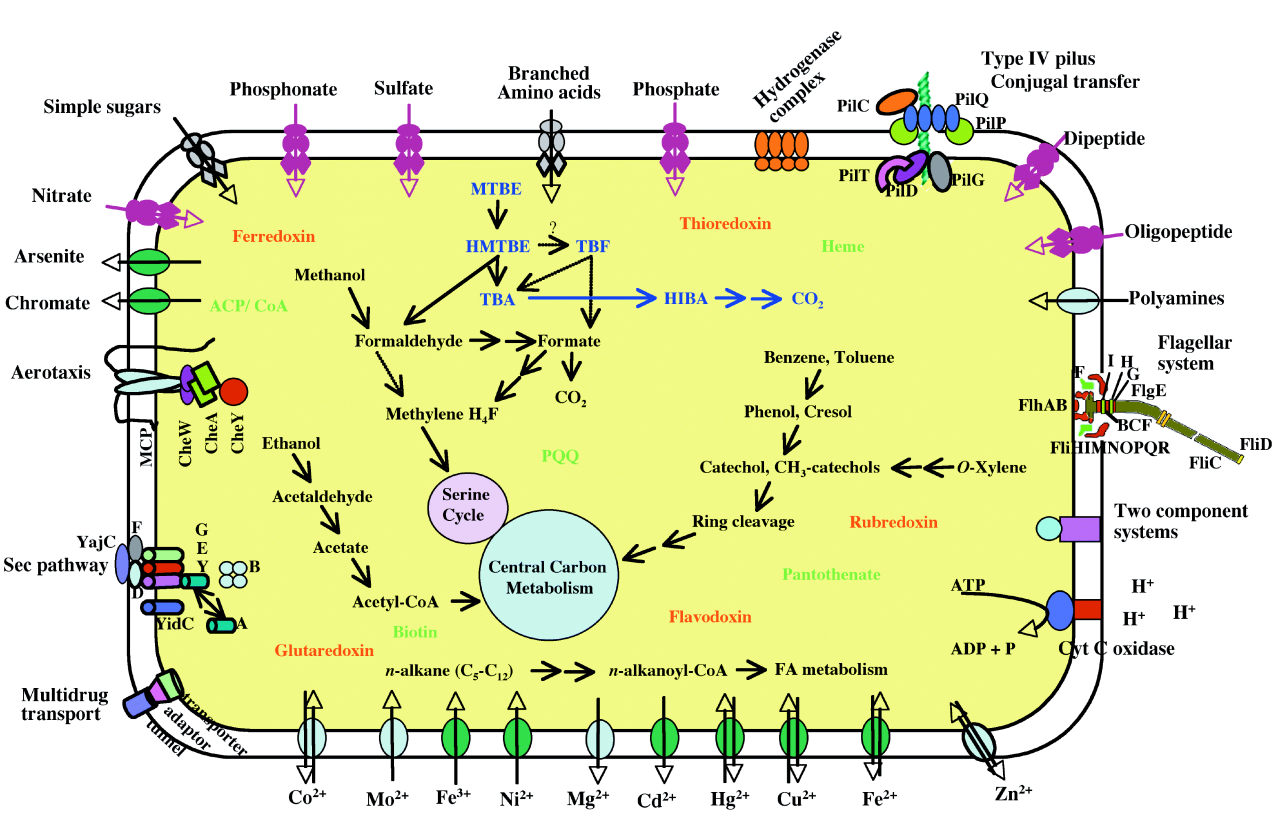
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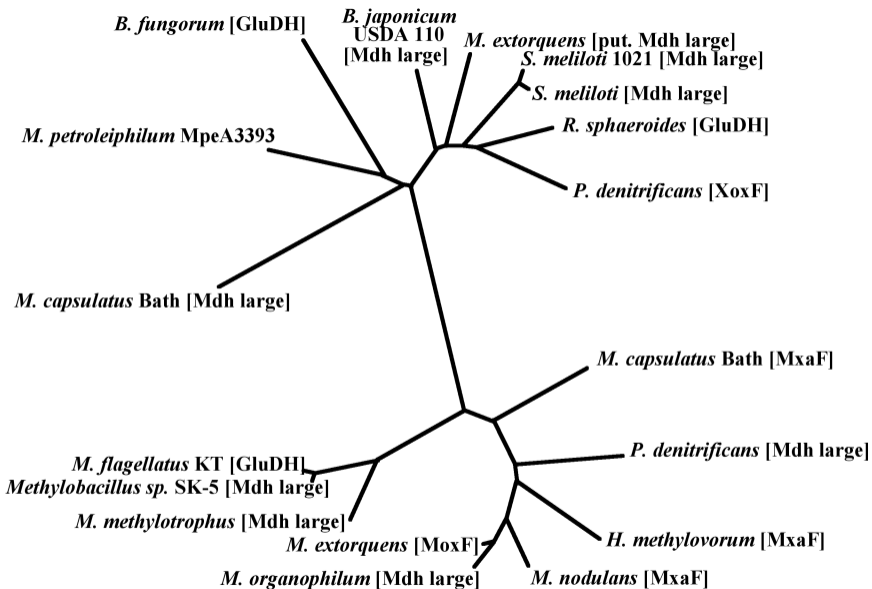


**B**

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